Welcome to STN International! Enter x:x

LOGINID:SSPTAEGS1646

PASSWORD:

SESSION RESUMED IN FILE 'BIOSIS, CAPLUS, EMBASE, MEDLINE' * * * * * * RECONNECTED TO STN INTERNATIONAL * * * * * TOTAL COPYRIGHT (C) 2007 AMERICAN CHEMICAL SOCIETY (ACS) FILE 'EMBASE' ENTERED AT 15:31:15 ON 27 JAN 2007 Copyright (c) 2007 Elsevier B.V. All rights reserved. FILE 'MEDLINE' ENTERED AT 15:31:15 ON 27 JAN 2007 Copyright (c) 2007 The Thomson Corporation FILE 'CAPLUS' ENTERED AT 15:31:15 ON 27 JAN 2007 SINCE FILE FILE 'BIOSIS' ENTERED AT 15:31:15 ON 27 JAN 2007 AT 15:31:15 ON 27 JAN 2007 COST IN U.S. DOLLARS

115.26 115.47 SESSION ENTRY FULL ESTIMATED COST

→ D Hist

(FILE 'HOME' ENTERED AT 14:40:59 ON 27 JAN 2007)

FILE 'BIOSIS, CAPLUS, EMBASE, MEDLINE' ENTERED AT 14:41:19 ON 27 JAN 2007

- 117 S ADIPOCYTOKINES AND PD<=20020726

 - 505 S ADIPONECTIN AND PD<=20020726 23383 S LEPTIN AND PD<=20020726
- 466840 S ("TYPE 2 DIABETES") OR OBESITY OR ("CARDIOVASCULAR
- 5288 S ("WHITE ADIPOSE TISSUE") AND PD<=20020726
- 0 S ("GLOBULAR DOMAIN" (XW) ADIPONECTIN) AND PD<=20020726 56 S LI AND L2

 - 46 DUP REM L7 (10 DUPLICATES REMOVED)
- 92 DUP REM L9 (69 DUPLICATES REMOVED) 161 S L2 AND L3
 - 60 S L 10 AND LA
- 60 DUP REM L11 (0 DUPLICATES REMOVED)
- IS DUP REM L13 (0 DUPLICATES REMOVED) 15 S L 10 AND L5
 - 128 S L2 AND L4
- 191 DUP REM L15 (137 DUPLICATES REMOVED)

⇒ D BIB ABS L12 4-25

L12 ANSWER 4 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:775209 CAPLUS <<LOGINID::20070127>>

- DN 138:37209
- TI Removal of visceral fat prevents insulin resistance and glucose intolerance of aging: an adipokine-mediated process?
- AU Gabriely, Ilan; Ma, Xiao Hui; Yang, Xiao Man; Atzmon, Gil; Rajala, Michael W.; Berg, Anders H.; Scherer, Phillip; Rossetti, Luciano; Barzilai, Nir
 - CS Diabetes Research and Training Center and Division of Endocrinology, Department of Medicine, Institute for Aging Research, Albert Einstein
 - College of Medicine, Bronx, NY, 10461, USA SO Diabetes (2002), 51(10), 2951-2958 CODEN: DIAEAZ; ISSN: 0012-1797
- PB American Diabetes Association
 - DT Journal
- LA English
- AB Age-dependent changes in insulin action and body fat distribution are risk factors for the development of type 2 diabetes
 - To examine whether the accumulation of visceral fat (VF) could play a direct role in the pathophysiol. of insulin resistance and type
 - 2 diabetes, we monitored insulin action, glucose
- tolerance, and the expression of adipo-derived peptides after surgical
- removal of VF in aging (20-mo-old) F344/Brown Norway (FBN) and in Zucker action were markedly impaired in aging FBN rats, and extraction of VF Diabetic Fatty (ZDF) rats. As expected, peripheral and hepatic insulin (accounting for .apprx. 18% of their total body fat) was sufficient to
- When examined at the mechanistic level, removal of VF in ZDF rats prevented restore peripheral and hepatic insulin action to the levels of young rats.
 - the progressive decrease in insulin action and delayed the onset of
 - diabetes, but VF extraction did not alter plasma free fatty acid levels
- leptin in s.c. (SC) adipose tissue were markedly decreased after However, the expression of tumor necrosis factor-a and
- VF removal (by approx. three- and twofold, resp.). Finally, extracted VF retained .apprx.15-fold higher resistin mRNA compared with SC fat. Our data suggest that insulin resistance and the development of diabetes can be significantly reduced in aging rats by preventing the age-dependent
- accumulation of VF. This study documents a cause-and-effect relationship between VF and major components of the metabolic syndrome.

RE.CNT 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L12 ANSWER 5 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 2002:824519 CAPLUS <<LOGINID::20070127>>
 DN 137:335596
 TI Significance of adipocytokine, fat-derived hormones, in metabolic syndrome AU Shimomura, lichiro; Funahashi, Tohru; Matsuzawa, Yuji

- CS Grad. Sch. Biofunct. Res., Osaka Univ., Japan SO Tanpakushitsu Kakusan Koso (2002), 47(14), 1896-1903 CODEN: TAKKAJ; ISSN: 0039-9450
 - PB Kyoritsu Shuppan
 - Journal; General Review П
 - LA Japanese
- metabolic syndromes including diabetes mellitus, hyperlipidemia, and adipocytokines, adipocyte-derived hormones, in obesity-caused A review on the pathophysiol. roles and clin. significance of atherosclerosis, focusing on PAI-1, TNF-a, leptin, and adiponectin. ΑB
- L12 ANSWER 6 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:768894 CAPLUS <<LOGINID::20070127>>
 - - 138:167458
- TI Syndrome of insulin resistance. Adipocytokines
- Dep. Internal Med. Molecular Sci., Grad. Sch. Med., Osaka Univ., Japan AU Kishida, Ken; Funahashi, Tohru CS Dep. Internal Med. Molecular Sei SO Saishin Igaku (2002), 57(8), 1799
 - Saishin Igaku (2002), 57(8), 1799-1805
- CODEN: SAIGAK; ISSN: 0370-8241
 - PB Saishin Igakusha
- DT Journal; General Review LA Japanese
- AB A review, on the roles of adipocytokines (fatty acids, glycerol, INFa, leptin, adiponectin) in the pathogenesis
 - of insulin resistance (IR) and IR-associated syndromes (obesity, type 2 diabetes, hypertension, lipid metabolic disorders, hyperinsulinemia, atherosclerosis)
- L12 ANSWER 7 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN
 - AN 2002:746437 CAPLUS << LOGINID::20070127>>
 - 138:150908
- Central role of adipocytokine on metabolic syndrome
- Dep. of Frontier Bioscience, Graduate School of Frontier Bioscience, Osaka AU Shimomura, lichiro; Funahashi, Tohru; Kihara, Shinji; Matsuzawa, Yuji University, Japan S
 - SO Jikken Igaku (2002), 20(12), 1762-1767
 - CODEN: JIIGEF; ISSN: 0288-5514
 - PB Yodosha
- DT Journal; General Review
- AB A review, on the roles of adipocytokines (PAI-I, TNF-a, LA Japanese

leptin, and adiponectin) on metabolic syndromes, such as obesity, diabetes, hyperlipemia, and atherosclerosis. L12 ANSWER 8 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN

AN 2002:469099 CAPLUS <<LOGINID::20070127>> DN 137:183818

TI Gene expression profile of rat adipose tissue at the onset of high-fat-diet obesity

AU Li, Jinping; Yu, Xinxin; Pan, Wentong; Unger, Roger H.

CS Gifford Laboratories, Touchstone Center for Diabetes Research, Department of Internal Medicine, University of Texas Southwestern Medical Center

Dallas, TX, 75390-8854, USA

SO American Journal of Physiology (2002), 282(6, Pt. 1),

E1334-E1341

CODEN: AJPHAP; ISSN: 0002-9513

PB American Physiological Society

DT Journal

adipose tissue (WAT) and requires recruitment of adipocyte precursor cells and their supporting infrastructure. To characterize the change in the LA English
AB Morbid obesity is the result of massive expansion of white

fat diet. Ninety-six genes were upregulated by at least 50% above the WAT the authors employed a cDNA subtraction screen for genes differentially expressed in epididymal fat pads harvested 1 wk after the start of a 60% , when adipocyte hypertrophy is present but hyperplasia is still minimal expression profile of the preexisting WAT at the start of obesity

leptin, adipocyte complement-related protein 30 kDa, and resistin, of control rats receiving a 4% fat diet. Of these genes, 30 had not previously been identified. Sixteen of the 96 genes, including

was a novel gene. Twenty-nine novel fragments were identified. Thus, at were predicted to encode a signal peptide. Ten of the 16 had been previously identified in other tissues and implicated in cell growth proliferation, differentiation, cell cycle control, and angiogenesis. the onset of high-fat-diet-induced obesity in rats, adipose

expansion of non-adipocyte tissues and of several uncharacterized novel factors. The only one of these thus far characterized functionally was tissue increases its expression of factors previously implicated in the

RE.CNT 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS found to promote lipogenesis. RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L12 ANSWER 9 OF 60 MEDLINE on STN
- AN 2002698566 MEDLINE <<LOGINID::20070127>> DN PubMed ID: 12388167
- TI Adiponectin is stimulated by adrenalectomy in ob/ob mice and is
 - highly correlated with resistin mRNA.
- AU Makimura Hideo; Mizuno Tooru M; Bergen Hugo; Mobbs Charles V

CS Neurobiology of Aging Laboratories, Fishberg Center for Neurobiology and Department of Geriatrics and Adult Development, Mount Sinai School of

Medicine, New York, New York 10029, USA

SO American journal of physiology. Endocrinology and metabolism, (2002 Dec) Vol. 283, No. 6, pp. E1266-71. Electronic Publication: 2002-08-13

ournal code: 100901226. ISSN: 0193-1849

United States

Journal; Article; (JOURNAL ARTICLE)

FS Priority Journals

English

EM 200212

ED Entered STN: 17 Dec 2002

Last Updated on STN: 5 Jan 2003 Entered Medline: 9 Dec 2002

plasma levels of resistin, an agent that some believe to confer insulin resistin expression were also highly correlated in diet-induced obese adrenalectomy can increase insulin sensitivity, we hypothesized that AB Plasma levels of the adipocyte product adiponectin, a putative Surprisingly, expression of adiponectin and resistin was highly positively correlated even after statistical removal of effects of adiponectin mRNA, adiponectin peptide, and resistin mRNA adrenalectomy would increase expression of adiponectin and adiponectin expression in ob/ob mice to wild-type levels and insulin, glucose, and adiposity. In addition, adiponectin and in adrenalectomized ob/ob mice. Adrenalectomy restored insulin-sensitizing agent, are reduced in obesity, whereas resistance, are thought to increase with obesity. Because stimulated adiponectin peptide to above wild-type levels. decrease expression of resistin. Therefore, we measured

L12 ANSWER 10 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:949636 CAPLUS <<LOGINID::20070127>>

mice. The data support a role for adiponectin in mediating some

effects of adrenalectomy on insulin sensitivity.

DN 138:220160

TI Resistin and adiponectin expression in visceral fat of obese rats: effect of weight loss

CS Endocrine-Metabolic Laboratory, Internal Medicine, Department of Medical AU Milan, Gabriella; Granzotto, Marnie; Scarda, Alessandro; Calcagno, Alessandra; Pagano, Claudio; Federspil, Giovanni; Vettor, Roberto

and Surgical Sciences, University of Padova, Padua, 35128, Italy

SO Obesity Research (2002), 10(11), 1095-1103

CODEN: OBREFR; ISSN: 1071-7323

PB North American Association for the Study of Obesity DT Journal

English

Obesity-related insulin resistance is closely associated with ₽ E

observed when s.c. adipose tissues of the same animals were compared. Weight resistin mRNA after weight loss does not support the hypothesis that resistin VAT of genetically obese in comparison with lean rats; no differences were whereas a further significant decrease in resistin mRNA level was observed we demonstrated the presence of resistin in immunocompetent cells in both reverse-transcription polymerase chain reaction. Moreover, we analyzed the immune system, thus suggesting an intriguing functional connection. Resistin is also present and equally expressed in splenocytes of lean and may play a causative role in insulin resistance in obese rats. Moreover, visceral fat accumulation. Several adipocyte-secreted mols. have been and resistin proteins. Some of these adipocytokines are also present in We determined adiponectin and resistin expressions in visceral (VAT) the variations after body-weight reduction in food-restricted obese rats levels after body-weight reduction, supporting its link with obesity related insulin resistance. On the contrary, the further decrease of humans and rats, thus adding another factor to the list of mols. that and s.c. adipose tissue of lean and obese Zucker (fa/fa) rats using obese rats. Adiponectin and resistin are down-regulated in VAT Resistin and adiponectin expression was significantly lower in loss resulted in an increase of adiponectin expression in VAT, of obese rats. Adiponectin expression is restored to normal diabetes, among them, the recently discovered adiponectin

adipose tissue shares with the immune system.

RE.CNT 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 11 OF 60 EMBASE COPYRIGHT (c) 2007 Elsevier B.V. All rights reserved on STN

AN 2002449010 EMBASE <<LOGINID::20070127>>

TI Cardiovascular risks in obesity.

AU Uchegbu E.C.; Kopelman P.G.

Wing, Royal London Hospital, Turner Street, London E1 1BB, United Kingdom. CS Dr. E.C. Uchegbu, Dept. of Diabetes and Metabolism, 5th Floor Alexandra

e.c.uchegbu@qmul.ac.uk SO Journal of Endocrinological Investigation, (2002) Vol. 25, No.

Refs: 38

10, pp. 915-918.

ISSN: 0391-4097 CODEN: JEIND7

Journal; General Review

CY Italy

Clinical Biochemistry FS 029

Cardiovascular Diseases and Cardiovascular Surgery 005

General Pathology and Pathological Anatomy

Endocrinology

- Public Health, Social Medicine and Epidemiology
 - LA English ED Entered STN: 3 Jan 2003

Last Updated on STN: 3 Jan 2003

DATA NOT AVAILABLE FOR THIS ACCESSION NUMBER

L12 ANSWER 12 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN

AN 2003:65783 CAPLUS <<LOGINID::20070127>>

TI Adipose tissue hormones DN 138:265785

AU Guerre-Millo, M.

CS Centre de Recherche des Cordeliers, Universite Pierre et Marie Curie,

Paris, 75006, Fr.

SO Journal of Endocrinological Investigation (2002), 25(10),

CODEN: JEIND7; ISSN: 0391-4097

PB Editrice Kurtis s.r.l. DT Journal; General Re

Journal; General Review LA English

AB A review. It is now widely accepted that white adipose tissue (WAT)

secretes a number of peptide hormones, including leptin, several angiotensinogen, plasminogen activator inhibitor-1 (PAI-1), cytokines, adipsin and acylation-stimulating protein (ASP),

This newly discovered secretory function has shifted the authors' view of adiponectin, resistin etc., and also produces steroids hormones.

energy homeostasis, glucose and lipid metabolism, vascular homeostasis, immune WAT, which is no longer considered only an energy storage tissue but a major endocrine organ, at the heart of a complex network influencing

response and even reproduction Virtually all known adipose secreted proteins are dysregulated when the WAT mass is markedly altered, either increased adiposesecreted products in the ethiopathol. and/or complications of both in the obese state or decreased in lipoatrophy. This strongly implicates obesity and cachexia. This review discusses the physiol.

hormones. Regulation of WAT secretion by the major regulatory factors relevance of adipose secretion by focusing on protein and steroid impinging on the adipocytes, i.e., insulin, glucocorticoids,

rationale for therapeutic strategies aimed at compensating adverse effects resulting from overprodn. or lack of a specific adipose secretory product catecholamines and thiazolidinediones (TZD) will be addressed. The

will be discussed.

RE.CNT 53 THERE ARE 53 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 13 OF 60 MEDLINE on STN AN 2002476461 MEDLINE <<LOGINID::20070127>>

- DN PubMed ID: 12238130
- II Glucose intolerance in visceral fat syndrome.
 - AU Matsuzawa Yuji
- Department of Internal Medicine and Molecular Science, Osaka University Graduate School.

SO Nippon rinsho. Japanese journal of clinical medicine, (2002 Jul)

Vol. 60 Suppl 7, pp. 746-51. Ref: 14 Journal code: 0420546. ISSN: 0047-1852.

CY Japan

DT Journal; Article; (JOURNAL ARTICLE) General Review; (REVIEW)

LA Japanese FS Priority Journals EM 200211

ED Entered STN: 20 Sep 2002

Last Updated on STN: 13 Dec 2002

Entered Medline: 20 Nov 2002

L12 ANSWER 14 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN

AN 2002:328227 CAPLUS <<LOGINID::20070127>> DN 137:335603

TI Obesity: Molecular mechanism of obesity and its complications

Shimomura, lichiro; Funahashi, Tohru; Matsuzawa, Yuji ΑU

Department of Internal Medicine and Molecular Science, Graduate School of Medicine, Osaka University, Japan S

SO Saishin Igaku (2002), 57(March, Zokango, Seikatsu Shukanbyo, Zenpen), 708-717

CODEN: SAIGAK; ISSN: 0370-8241

DT Journal; General Review PB Saishin Igakusha

adiponectin, and resistin) that are related to lipid metabolism and LA Japanese
AB A review on mol. factors (especially PAI-1, TNF-a, leptin,

visceral fat accumulation in human.

L12 ANSWER 15 OF 60 EMBASE COPYRIGHT (c) 2007 Elsevier B.V. All rights reserved on STN

TI Adiponectin: A link between excess adiposity and associated AN 2002422400 EMBASE <<LOGINID::20070127>>

comorbidities?

AU Ukkola O.; Santaniemi M.

CS O. Ukkola, Department of Internal Medicine, Biocenter Oulu, University of

Oulu, Kajaaninie 50, 90220 Oulu, Finland. olavi.ukkola@oulu.fi SO Journal of Molecular Medicine, (2002) Vol. 80, No. 11, pp.

696-702.

ISSN: 0946-2716 CODEN: JMLME8 Journal; General Review Germany ჯ

Endocrinology ES

Internal Medicine

Drug Literature Index

LA English SL English

Last Updated on STN: 5 Dec 2002 ED Entered STN: 5 Dec 2002

adipose tissue. In contrast to other adipocy-tokines, adiponectin AB Adiponectin is a novel polypeptide that is highly specific to evels are decreased in obesity and associated comorbidities, such as type 2 diabetes. Decreased

It has been suggested that several agents, such as tumor necrosis factor expression of adiponectin is correlated with insulin resistance.

mechanisms for the development of atherosclerotic vascular disease in a, could mediate their effects on insulin metabolism through modulating adiponectin secretion from adipocytes. The

bese individuals are largely unknown. Several findings support the nteresting hypothesis that adiponectin could be a link between obesity and related atherosclerosis. First, adiponectin

adiponectin is accumulated more preferably to the injured vascular levels are lower in patients with coronary artery disease. Second, adiponectin modulates endothelial function and has an inhibitory effect on vascular smooth muscle cell proliferation. Moreover,

wall than intact vessels and has been shown to suppress macrophage-to-foam modulation of inflammation. Thiazolidinediones, antiatherogenic and other cell transformation. Adiponectin may also be involved in the

effects have been explained by their direct enhancing effect on adiponectin. In conclusion, adiponectin has

beneficial effects on metabolism. Therefore it is not a surprise that anti-inflammatory and anti-atherogeneic effects as well as multiple obesity, and it has been shown to ameliorate hyperglycemia and adiponectin therapy has been tested in animal models of

hyperinsulinemia without inducing weight gain or even inducing weight loss

evidence of an association between adiponectin and the metabolic in some studies. Unlike agents that exert their effects centrally, and cardiovascular complications of obesity is growing all the adiponectin's effects seem to be peripherally mediated. The

L12 ANSWER 16 OF 60 MEDLINE on STN AN 2002670544 MEDLINE <<LOGNID::20070127>>

DN PubMed ID: 12430302

TI Glucose metabolism in adipose tissue.

AU Inoue Atsushi; Tobe Kazuyuki; Suzuki Ryo; Kadowaki Takashi

CS Department of Internal Medicine, Graduate School of Medicine, University of Tokyo.

SO Nippon rinsho. Japanese journal of clinical medicine, (2002 Oct) Vol. 60 Suppl 10, pp. 673-80. Ref: 21

Journal code: 0420546. ISSN: 0047-1852.

DT Journal; Article; (JOURNAL ARTICLE) CY Japan

General Review; (REVIEW)

LA Japanese FS Priority Journals EM 200302

ED Entered STN: 15 Nov 2002

Last Updated on STN: 21 Feb 2003

Entered Medline: 20 Feb 2003

L12 ANSWER 17 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN

AN 2003:262668 CAPLUS << LOGINID:: 20070127>>

DN 138:399708

variables of fibrinolysis in overweight and obese hypertensive patients TI Relationship between IL-6, leptin and adiponectin and

AU Skurk, T.; van Harmelen, V.; Lee, Y. -M.; Wirth, A.; Hauner, H.

German Diabetes Research Institute at the Heinrich-Heine-University, S

Dusseldorf, 40225, Germany

SO Hormone and Metabolic Research (2002), 34(11/12), 659-663

CODEN: HMMRA2; ISSN: 0018-5043

PB Georg Thieme Verlag DT Journal

LA English
AB Impaired fibrinolysis is a common finding in obese humans. This condition is now considered as an established risk factor for thromboembolic

specific pattern of circulating concns. of fat-cell products interleukin-6 complications. Furthermore, obesity is characterized by a

was to investigate the relationship between these proteins and selected (IL-6), leptin, and adiponectin. The aim of our study

variables of the fibrinolytic system in 74 mildly hypertensive, overweight subjects. Circulating IL-6 and leptin levels showed a pos.

association with BMI (r = 0.24, p = 0.04 and r = 0.70, p < 0.0001), whereas adiponectin was not correlated to BMI. Interestingly, IL-6 was

also pos. associated with t-PA/PAI-1 complexes after adjustment for BMI and other anthropometric variables. Leptin was pos. correlated with

PAI-1 activity and antigen (r = 0.32, p = 0.006) and r = 0.37, p < 0.001,

assocns. lost significance after correction for BMI or HOMA, an insulin resp.) and neg. with t-PA activity (r = -0.27, p = 0.03). However, these

ndependently and neg. correlated with PAI-1 antigen (r = -0.26, p = 0.04, after correction for BMD. In conclusion, our study provides further sensitivity index. In contrast, adiponectin levels were evidence that IL-6, leptin, and adiponectin are

RE.CNT 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS associated with impaired fibrinolysis in overweight hypertensive humans. RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 18 OF 60 EMBASE COPYRIGHT (c) 2007 Elsevier B.V. All rights reserved on STN

AN 2005364778 EMBASE <<LOGINID::20070127>>

TI Adipose tissue as an endocrine organ

AU Prins J.B.

Dr. J.B. Prins, Princess Alexandra Hospital, Ipswich Rd, Woolloongabba, QLD 4102, Australia S

SO Best Practice and Research in Clinical Endocrinology and Metabolism, (2002) Vol. 16, No. 4, pp. 639-651...

Refs: 97

ISSN: 1521-690X CODEN: BPRCE

CY United Kingdom

DT Journal; General Review 903 S

Clinical Biochemistry Endocrinology 620

LA English

SL English Э

Last Updated on STN: 27 Oct 2005 Entered STN: 27 Oct 2005

nighlighted, and areas in need of future research are suggested. .COPYRGT. soluble products with both local and distant actions. These hormones have endocrinology of adipose tissue by concentrating on functional aspects of mportant roles in metabolism, reproduction, cardiovascular function and AB Adipose tissue is a highly active endocrine organ secreting a range of influences other organ systems, including the brain, liver and skeletal regulated by nutritional status, and both are inextricably linked to the the secreted products. The data of particular relevance to humans are mmunity. It is now evident that adipose endocrine function directly muscle. The endocrine function of adipose tissue is significantly energy storage role of adipose tissue. This chapter highlights the 2002 Elsevier Science Ltd. All rights reserved.

L12 ANSWER 19 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN

2003:262661 CAPLUS << LOGINID::20070127>>

AU Atzmon, G.; Yang, X. M.; Muzumdar, R.; Ma, X. H.; Gabriely, I.; Barzilai DN 138:396443 TI Differential gene expression between visceral and subcutaneous fat depots

Department of Medicine, Albert Einstein College of Medicine, Bronx, NY Institute for Aging Research & Diabetes Research and Training Center 10461, USA

SO Hormone and Metabolic Research (2002), 34(11/12), 622-628 CODEN: HMMRA2; ISSN: 0018-5043

PB Georg Thieme Verlag

DT Journal

LA English
AB Abdominal obesity has been linked to the development of insulin resistance and Type 2 diabetes mellitus

fat from 5 young rats. We extracted mRNA from the fat tissue and performed gene array hybridization using Affymetrix technol. with a platform containing 9000 genes. Out of the 1660 genes that were expressed in fat tissue, 297 (17.9%) genes show a two-fold or higher difference in their expression (DM2). By surgical removal of visceral fat (VF) in a variety of rodent peripheral fat depots, we obtained perirenal visceral (VF) and s.c. (SC) establishing a cause-effect relationship between VF and the metabolic syndrome. To characterize the biol. differences between visceral and models, we prevented insulin resistance and glucose intolerance,

glucose homeostasis, insulin action, and lipid metabolism We confirmed the higher in VF fat (by 3-7 fold) and the 20 genes whose expression is higher in SC fat (by 3-150 fold), many of which are predominantly involved in between the two tissues. We present the 20 genes whose expression is

findings of gene array expression and quantified the changes in expression in VF of genes involved in insulin resistance (PPARg leptin

demonstrated increased expression of resistin in VF by around 12-fold and inhibitor-1, PAI-1) by real-time PCR (qRT-PCR) technol. Finally, we gene expression platform. These results indicate that visceral fat and) and its syndrome (angiotensinogen and plasminogen activating adiponectin by around 4-fold, peptides that were not part of the

s.c. fat are biol. distinct.
RE.CNT 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 20 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation 등

STS

AN 2002:387248 BIOSIS <<LOGINID::20070127>>

ON PREV200200387248

Resistin and adiponectin expression in lean and obese Zucker

Nutrition Sciences, University of Alabama at Birmingham, 1675 University AU Blaylock, Matthew L. [Reprint author]; Nagy, Tim R. [Reprint author] Blvd, Birmingham, AL, 35294, USA S

- SO FASEB Journal, (March 20, 2002) Vol. 16, No. 4, pp. A603. print. Meeting Info.: Annual Meeting of the Professional Research Scientists on Experimental Biology. New Orleans, Louisiana, USA. April 20-24, 2002 CODEN: FAJOEC, ISSN: 0892-6638.
 - DT Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

ED Entered STN: 17 Jul 2002 LA English

AB The mechanisms underlying obesity and type 2 Last Updated on STN: 17 Jul 2002

rats over a range of ages. Animals (n=9-10/group) were euthanized at 6, 7, 10, and 14 weeks of age and epididymal white adipose tissue was more, had greater adipose tissue mass as well as higher levels of plasma was reduced in the fatty compared to the lean Zucker rats (p<0.05). Our results are in agreement with recently published data suggesting that the collected. The results showed that the fatty rats weighed significantly Within each age class, the expression of resistin and adiponectin these processes. The purpose of this study was to determine the expression of resistin and adiponectin in lean and fatty Zucker diabetes remain to be elucidated. Two novel adipose-derived cytokines, resistin and adiponectin, have been implicated in leptin, insulin, free-fatty acids, and triglycerides (p<0.05). expression of resistin and adiponectin is reduced with obesity and increasing insulin resistance.

- L12 ANSWER 21 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN 2002:424393 CAPLUS << LOGINID:: 20070127>>

 - DN 138:2729
- TI A novel transgenic mouse model of visceral fat obesity and
- Medicine, Harvard Medical School and Beth Israel Deaconess Medical Center, CS Division of Endocrinology, Diabetes and Metabolism, Department of AU Masuzaki, Hiroaki; Flier, Jeffrey S. metabolic syndrome 3oston, MA, USA
 - SO Molecular Medicine (Tokyo, Japan) (2002), 39(4), 464-474 CODEN: MOLMEL; ISSN: 0918-6557
 - PB Nakayama Shoten

 - Journal; General Review Japanese
- decreased energy metabolism, glucose tolerance, and insulin sensitivity in aP2 metabolic disorders; (2) transgenic mouse (aP2 HSD-1 mouse) model of ncreased expression of lipoprotein lipase, angiotensinogen, and tumor AB A review. The topics discussed are (1) visceral fat obesity and visceral fat obesity by overexpressing adipose tissue specific 11b-hydroxysteroid dehydrogenase type 1 (11b-HDS-1); (3) HSD-1 mice; (4) leptin resistance, visceral obesity,

- (PEPCK), and glucose-6-phosphatase (G6Pase) in liver of aP2 HSD-1 mice. and resistin in adipose tissues of aP2 HSD-1 mice; and (5) increases in free fatty acids, corticosterone, phosphoenolypyruvate carboxykinase
- L12 ANSWER 22 OF 60 EMBASE COPYRIGHT (c) 2007 Elsevier B.V. All rights reserved on STN
 - AN 2003447834 EMBASE <<LOGINID::20070127>>
 - II Slimming down without DGAT.
 - AU Brazil M.
- SO Nature Reviews Drug Discovery, (2002) Vol. 1, No. 6, pp. 408. Refs: 1

- ISSN: 1474-1776 CODEN: NRDDAG
 - United Kingdom
 - Journal; Note 8 DT
- Endocrinology
- Clinical Biochemistry 030 029
- Drug Literature Index Pharmacology 4 6 1 037

 - LA English
- Last Updated on STN: 20 Nov 2003 ED Entered STN: 20 Nov 2003
- DATA NOT AVAILABLE FOR THIS ACCESSION NUMBER
- L12 ANSWER 23 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN
 - AN 2003:16789 CAPLUS <<LOGINID::20070127>> DN 138:151453
- TI The morbid condition caused by insulin resistance in the obese patient with diabetes. Homology of leptin, adiponectin and LPL
- AU Takahashi, Toshikazu; Mochihara, Yuki; Kodate, Shinya; Mashimo, Ikuo;
- Maehata, Eisuke; Shiba, Teruo; Yamakado, Minoru; Inoue, Minoru; Taniyama, CS Quality Assurance Office, Surnikin Bioscience K. K., Sagamihara, 229-1125, Tazawa, Hiromitsu; Taira, Yoshihisa; Yano, Masao; Shimomura, Koji Matsuo; Suzuki, Seiji
- Seibutsu Shiryo Bunseki (2002), 25(5), 385-391 S
 - CODEN: SSBUEL; ISSN: 0913-3763 PB Seibutsu Shiryo Bunseki Kagakkai
- Journal
 - LA Japanese
- AB We compared the decrease in insulin resistivity in obese patients with diabetes using the insulin resistance index (HOMA-R) and the physiol. adiponectin). Changes in the homologous levels were investigated active substances secreted by adipose tissue (LPL mass, leptin, at the same time. As the HOMA-R progressed, leptin became

which HOMA-R was over 2.0. From these results, insulin resistivity showed <0.0022) were recognized, and the same when compared with the group in relationship. As for the adipocytokine correlation, in the group with a 0.694, p = 0.0003) and adiponectin vs. LPL mass (r = 0.618, p BMI exceeding 25 kg/m2, correlations of BMI vs. leptin (r = tinetic and adiponectin increased, showing an antagonistic characteristic morbid condition.

- L12 ANSWER 24 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN
 - 2002:587011 CAPLUS << LOGINID:: 20070127>>
 - 137:382874
- TI Increased fat intake, impaired fat oxidation, and failure of fat cell proliferation result in ectopic fat storage, insulin resistance, and ype 2 diabetes mellitus
- Ravussin, Eric; Smith, Steven R.
- Pennington Biomedical Research Center, Baton Rouge, LA, 70808-4124, USA SS
 - Annals of the New York Academy of Sciences (2002), 967(Lipids
 - and Insulin Resistance), 363-378
- CODEN: ANYAA9; ISSN: 0077-8923
 - PB New York Academy of Sciences
 - Journal; General Review D
 - English
- A review. It is widely accepted that increasing adiposity is associated with insulin resistance and increased risk of type 2

hypothesis. As such, two new paradigms have emerged that may explain the nd insulin resistance. The endocrine paradigm developed in parallel with support the acquired lipodystrophy hypothesis as a link between adiposity he failure of the adipose tissue mass to expand and thus to accommodate established links between adiposity and disease. Three lines of evidence probably the beta cell. The importance of this finding is exemplified by the portal/visceral hypothesis. This hypothesis proposes that increased ipodystrophy, results in severe insulin resistance and diabetes. This is Third, increased fat cell size is highly associated with insulin resistance adequate adipose tissue mass in either mice or humans, also known as and the development of diabetes. Increased fat cell size may represent an increased energy influx. Taken together, these three observations several studies demonstrating that the degree of lipid infiltration into obese patients also shunt lipid into the skeletal muscle, the liver, and muscle, and the pancreatic insulin-secreting beta cell. Second, most adiposity, particularly in the visceral depots, leads to increased free fatty acid flux and inhibition of insulin action via Randle's effect in hought to be the result of ectopic storage of lipid into liver, skeletal support the ectopic fat storage syndrome. First, failure to develop skeletal muscle and liver correlates highly with insulin resistance. insulin-sensitive tissues. Recent data do not entirely support this diabetes. The predominant paradigm used to explain this link is

metabolism of distant tissues. These two new paradigms provide a framework to advance our understanding of the pathophysiol. of the insulin-resistance the ectopic fat storage syndrome hypothesis. Adipose tissue secretes a endocrine gland, secreting numerous factors with potent effects on the resistin. From this viewpoint, adipose tissue plays a critical role as an angiotensin II, adiponectin (also called ACRP30 and adipoQ), and variety of endocrine hormones, such as leptin, interleukin-6, syndrome.

RE.CNT 118 THERE ARE 118 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L12 ANSWER 25 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN
- 2002:907355 CAPLUS <<LOGINID::20070127>> Ş
 - DN 138:300852
- An adipocentric view of signaling and intracellular trafficking
 - AU Mora, Silvia; Pessin, Jeffrey E.
- Department of Physiology and Biophysics, The University of Iowa, Iowa, IA, USA S
 - SO Diabetes/Metabolism Research and Reviews (2002), 18(5), 345-356
 - CODEN: DMRRFM; ISSN: 1520-7552 PB John Wiley & Sons Ltd.
 - DT Journal; General Review

 - LA English
- AB A review. Adipocytes have traditionally been considered to be the primary stimulate both glucose uptake and lipogenesis. Conventional wisdom held site for whole body energy storage mainly in the form of triglycerides and models and humans also leads to metabolic disorders that result in severe opposite functions can be resolved by the establishment of adipocytes not regulating insulin secretion, insulin action, glucose and lipid metabolism, secretory factors include enzymes (lipoprotein lipase (LPL) and adipsin) increased adipose tissue mass and/or increased lipolysis and circulating mol. function of these adipocyte-derived signals are poorly understood, growth factors [vascular endothelial growth factor (VEGF)], cytokines secretes a variety of signaling mols. into the circulation. Although the resistance and perhaps diabetes. However, it has become increasingly only as a fuel storage depot but also as a critical endocrine organ that energy balance, host defense and reproduction The diversity of these they play a central role in the maintenance of energy homeostasis by states of insulin resistance and potential diabetes. These apparently apparent that loss of adipose tissue (lipodystrophies) in both animal tumor necrosis factor-a, interleukin 6) and several other hormones fatty acids. This occurs through the ability of insulin to markedly that defects in fuel partitioning into adipocytes either because of involved in fatty acid and glucose metabolism (leptin, Acrp30, free fatty acids resulted in dyslipidemia, obesity, insulin

the GLUT4 glucose transporter.

RE.CNT 174 THERE ARE 174 CITED REFERENCES AVAILABLE FOR THIS and on one of the most intensively studied regulated membrane proteins, resistin and acylation stimulation protein). Despite the large number of current knowledge of the trafficking and secretion processes that take mols. secreted by adipocytes, our understanding of the pathways and adipocytes is poorly understood. In this article, we will review the mechanisms controlling intracellular trafficking and exocytosis in place in adipocytes, focusing our attention on two of the best characterized adipokine mols. (leptin and adiponectin) RECORD

VEL CITATIONS AVAILABLE IN THE RE FORMAT

=> D bib ABS 112 33-35, 41, 45, 49, 50-53, 55-60

L12 ANSWER 33 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:786585 CAPLUS << LOGINID::20070127>>

DN 138:83795

in white and brown adipose tissues: influence of b3-adrenergic TI Regulation of adiponectin and leptin gene expression agonists, retinoic acid, leptin and fasting

Zhang, Yi; Matheny, Michael; Zolotukhin, Sergei; Turner, Nihal; Scarpace, Philip J. ΥC

CS Department of Veterans Affairs Medical Center, Geriatric Research, Education and Clinical Center, Gainesville, FL, 32608-1197, USA

SO Biochimica et Biophysica Acta, Molecular and Cell Biology of Lipids (

CODEN: BBMLFG; ISSN: 1388-1981 2002), 1584(2-3), 115-122

PB Elsevier B.V.

Journal

English

two adipocyte-derived hormones may be simultaneously influenced by common obesity-related factors. The authors examined adiponectin mRNA levels in WAT and in some instances, brown adipose tissue (BAT) ollowing fasting and refeeding, acute and chronic administration of a reported for most of the treatments. Fasting diminished and refeeding 53-adrenergic agonist, acute treatment with retinoic acid (RA) and a in each circumstance. Serum concns. of adiponectin were also eripheral injection of the b3-adrenergic agonist, CL316,243, glucocorticoid, and following chronic infusion of leptin and evels rise with obesity, suggesting that regulation of these compared the expression of adiponectin to that of leptin reversed both adiponectin and leptin gene expression. Circulating adiponectin levels fall, whereas leptin

suppressed both leptin and adiponectin expression in

BAT was also observed following this treatment. Although CL316,23 lowered serum concns. in contrast to suppressions in both mRNA and serum levels of short-term regulation of the two adipokine expression in WAT is somewhat synthesis in WAT compared to controls, but prevented the reduction in similar, perhaps subjective to common control of energy balance. The WAT. A small but significant reduction in adiponectin expression in adiponectin synthesis associated with pair feeding. Food restriction long-term regulation of adiponectin expression in WAT appears to through pair feeding also diminished adiponectin expression in resulted in an elevation of adiponectin expression in WAT and be the opposite of that of leptin and may be more sensitive to leptin by a similar treatment as previously reported. Chronic adiponectin levels. A chronic 7-day infusion of CL316,243 appear to participate directly in adiponectin synthesis. The are inversely correlated with obesity, leptin does not serum leptin levels markedly, it did not affect serum BAT. Collectively, although leptin and adiponectin administration of leptin did not alter adiponectin

changes in adiposity or insulin sensitivity.

RE.CNT 35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS. RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 34 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN AN 2003:785040 CAPLUS <<LOGINID::20070127>> Z Z

140:233198

Adiponectin and resistin

Graduate School of Medicine, Osaka University, Japan

 TI Adiponectin and resistin
 AU Ohuchi, Noriari; Funabashi, Toru; Matsuzawa, Yuji
 CS Graduate School of Medicine, Osaka University, Jap.
 SO Bunshi Tonyobyogaku no Shinpo (2002) 53-59 Bunshi Tonyobyogaku no Shinpo (2002) 53-59

CODEN: BTSHFO

Kanehara Shuppan В

Journal; General Review

atherogenesis suppression and improved insulin sensitivity; (3) resistin adipocyte-derived plasma protein adiponectin and its effects on (4) other adipocytokines tumor necrosis factor-a (TNF-a) and LA Japanese
AB A review. The topics discussed are (1) adipocytokines; (2) expression in relation to obesity and insulin resistance; and

L12 ANSWER 35 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

STS

AN 2002:374089 BIOSIS <<LOGINID::20070127>> DN PREV200200374089

- TI Control of energy homeostasis and insulin action by adipocyte hormones: Leptin, acylation stimulating protein, and adiponectin.
 - Havel, Peter J. [Reprint author] ΑU
- CS Department of Nutrition, University of California, Davis, One Shields Avenue, Davis, CA, 95616, USA
- pjhavel@ucdavis.edu
- SO Current Opinion in Lipidology, (February, 2002) Vol. 13, No. 1,
 - pp. 51-59. print. ISSN: 0957-9672.

 - DT Article
- General Review; (Literature Review) English ځ
 - Entered STN: 3 Jul 2002 田
 - Last Updated on STN: 3 Jul 2002
- L12 ANSWER 41 OF 60 EMBASE COPYRIGHT (c) 2007 Elsevier B.V. All rights reserved on STN
 - AN 2002317642 EMBASE <<LOGINID::20070127>>
- Adiponectin enhances insulin action by decreasing ectopic fat
- deposition.
- Ravussin E. ΥU
- Enhancement Ctr., 6400 Perkins Rd., Baton Rouge, LA 70808 4124, United E. Ravussin, Pennington Biomedical Research Ctr., Health/Performance States. ravusse@pbrc.edu ပ္ပ
 - SO Pharmacogenomics Journal, (2002) Vol. 2, No. 1, pp. 4-7.
 - Refs: 16
- ISSN: 1470-269X CODEN: PJHOAZ CY United Kingdom
 - Journal; Article
- Endocrinology
- Human Genetics
 - Pharmacology 4 6 1
- Drug Literature Index

LA English

- ED Entered STN: 19 Sep 2002
- Last Updated on STN: 19 Sep 2002 DATA NOT AVAILABLE FOR THIS ACCESSION NUMBER
- L12 ANSWER 45 OF 60 MEDLINE on STN AN 2001666524 MEDLINE <<LOGINID::20070127>> N N
- TI The molecular mechanisms by which PPAR gamma/RXR inhibitors improve PubMed ID: 11712415 insulin resistance.
- CS Department of Metabolic Diseases, Graduate School of Medicine, University AU Yamauchi T; Kadowaki T

- SO Nippon rinsho. Japanese journal of clinical medicine, (2001 Nov) Vol. 59, No. 11, pp. 2245-54. Ref: 22 Journal code: 0420546. ISSN: 0047-1852.
- Journal; Article; (JOURNAL ARTICLE) CY Japan DT Journal
 - General Review; (REVIEW)
 - LA Japanese
- FS Priority Journals EM 200201
- ED Entered STN: 20 Nov 2001
- Last Updated on STN: 28 Jan 2002

Entered Medline: 25 Jan 2002

- AB Potent activation of PPAR gamma by thiazolidinediones(TZD) increases TG content of WAT, thereby decreasing TG content of liver/muscle, leading to
 - Moderate reduction of PPAR gamma activity by PPAR gamma/RXR inhibitors amelioration of insulin resistance at the expense of obesity.
- and increase in fatty-acid combustion and decrease in lipogenesis, thereby ameliorating HF diet-induced obesity and insulin resistance.

decreases TG content of WAT/muscle/liver due to increased leptin

- Moreover, PPAR gamma/RXR inhibitors decrease lipogenesis in WAT, while TZD stimulate adipocyte differentiation and apoptosis, thereby both preventing
 - adipocyte hypertrophy, which is associated with alleviation of insulin resistance presumably due to decreases in FFA, and TNF alpha, and upregulation of adiponectin. We conclude that although by
- different mechanisms, both PPAR gamma/RXR inhibitors and PPAR gamma agonist improve insulin resistance, which is associated with decreased TG content of muscle/liver and prevention of adipocyte hypertrophy.
- L12 ANSWER 49 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation
- AN 2001:397010 BIOSIS <<LOGINID::20070127>> DN PREV200100397010
- TI The fat-derived hormone adiponectin reverses insulin resistance associated with both lipoatrophy and obesity.
- AU Yamauchi, T.; Kamon, J.; Waki, H.; Terauchi, Y.; Kubota, N.; Hara, K.;
- Shudo, K.; Yoda, M.; Nakano, Y.; Tobe, K.; Nagai, R.; Kimura, S.; Tomita, Mori, Y.; Ide, T.; Murakami, K.; Tsuboyama-Kasaoka, N.; Ezaki, O.; Akanuma, Y.; Gavrilova, O.; Vinson, C.; Reitman, M. L.; Kagechika, H.;
- CS Department of Internal Medicine, Graduate School of Medicine, University M.; Froguel, P.; Kadowaki, T. [Reprint author] of Tokyo, Tokyo, Japan

 - kadowaki-3im@h.u-tokyo.ac.jp SO Nature Medicine, (August, 2001) Vol. 7, No. 8, pp. 941-946.
- ISSN: 1078-8956

- Arricle
- English
- Last Updated on STN: 22 Feb 2002 ED Entered STN: 22 Aug 2001
- molecules involved in both fatty-acid combustion and energy dissipation in muscle. Moreover, insulin resistance in lipoatrophic mice was completely is implicated in the development of insulin resistance in mouse models of decreases insulin resistance by decreasing triglyceride content in muscle and liver in obese mice. This effect results from increased expression of diabetes and metabolic syndrome to chromosome 3q27, where the gene AB Adiponectin is an adipocyte-derived hormone. Recent genome-wide reversed by the combination of physiological doses of adiponectin encoding adiponectin is located. Here we show that decreased both obesity and lipoatrophy. These data also indicate that the expression of adiponectin correlates with insulin resistance in replenishment of adiponectin might provide a novel treatment or leptin alone. We conclude that decreased adiponectin mouse models of altered insulin sensitivity. Adiponectin scans have mapped a susceptibility locus for type 2 and leptin, but only partially by either adiponectin modality for insulin resistance and type 2
- L12 ANSWER 50 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN
 - 2002:34308 CAPLUS << LOGINID:: 20070127>>
 - 137:150289
- Insulin resistance and cytokines

AU Hirose, Hiroshi; Yajima, Ken; Yamamoto, Hiroyuki; Kawai, Toshihide; Ishii, Kanako; Hayashi, Keisuke; Kawabe, Hiroshi; Saito, Ikuo; Saruta, Takao Tatsuya; Fujita, Haruhisa; Seto, Yoshiko; Miyashita, Kiichi; Nishikai,

- CS School of Medicine, Department of Internal Medicine, Keio University,
- Diabetes Frontier (2001), 12(5), 590-596 ၀ွ
 - CODEN: DIFREZ; ISSN: 0915-6593
 - PB Medikaru Rebyusha
- Journal; General Review Japanese
- liabetes, hypertension, and hyperlipidemia. Adipocytokines discussed are AB A review on contributions of adipocytokines adipose-derived bioactive substances in the development of insulin resistance in obesity, free fatty acids, tumor necrosis factor-a, leptin, adiponectin, and resistin.
- L12 ANSWER 51 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

- AN 2001:540867 BIOSIS <<LOGINID::20070127>> DN PREV200100540867
- TI Physiological role of adipose tissue: White adipose tissue as an endocrine and secretory organ.
 - AU Trayhum, Paul [Reprint author]; Beattie, John H.
- CS Department of Medicine, University Clinical Departments, University of Liverpool, Liverpool, L69 3GA, UK
 - SO Proceedings of the Nutrition Society, (August, 2001) Vol. 60. p_trayhum@altavista.com
 - No. 3, pp. 329-339. print. CODEN: PNUSA4. ISSN: 0029-6651.
 - - Article Ы
- General Review; (Literature Review) LA English

 - ED Entered STN: 21 Nov 2001
 - Last Updated on STN: 25 Feb 2002
- normal glucose homeostasis and a role in inflammatory processes has been AB The traditional role attributed to white adipose tissue is energy storage fatty acids being released when fuel is required. The metabolic role of white fat is, however, complex. For example, the tissue is needed for proposed. A radical change in perspective followed the discovery of principally by white fat, giving the tissue an endocrine function. leptin; this critical hormone in energy balance is produced
 - adipocytes, which include angiotensinogen, adipsin, acylation-stimulating protein, adiponectin, retinol-binding protein, tumour neorosis Leptin is one of a number of proteins secreted from white
- haemostasis or the complement system. The effects of specific proteins factor. Some of these proteins are inflammatory cytokines, some play a factor alpha, interleukin 6, plasminogen activator inhibitor-1 and tissue role in lipid metabolism, while others are involved in vascular
 - adipose tissue. The most recently described adipocyte secretory proteins may be autocrine or paracrine, or the site of action may be distant from are fasting-induced adipose factor, a fibrinogen-angiopoietin-related tissue-specific factor which is reported to induce insulin resistance, protein, metallothionein and resistin. Resistin is an adipose
- and stress-response protein which may have an antioxidant role. The key each secreted protein, and to assess the pathophysiological consequences of changes in adipocyte protein production with alterations in adiposity identify the complement of secreted proteins, to establish the role of challenges in establishing the secretory functions of white fat are to linking diabetes to obesity. Metallothionein is a metal-binding
- evidence of links between increased production of some adipocyte factors In essence, white adipose tissue is a major secretory and endocrine organ involved in a range of functions beyond simple fat storage. obesity, fasting, cachexia). There is already considerable

L12 ANSWER 56 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

AN 2000:304930 BIOSIS <<LOGINID::20070127>> DN PREV200000304930

Tl Plasma concentrations of a novel, adipose-specific protein,

adiponectin, in type 2 diabetic patients.

AU Hotta, Kikuko [Reprint author]; Funahashi, Tohru; Arita, Yukio; Takahashi, Muraguchi, Masahiro; Ohmoto, Yasukazu; Nakamura, Tadashi; Yamashita Kuriyama, Hiroshi; Ouchi, Noriyuki; Maeda, Kazuhisa; Nishida, Makoto; Kihara, Shinji; Sakai, Naohiko; Nakajima, Tadahisa; Hasegawa, Kyoichi; Masahiko; Matsuda, Morihiro; Okamoto, Yoshihisa; Iwahashi, Hiromi; Shizuya; Hanafusa, Toshiaki; Matsuzawa, Yuji

CS Department of Internal Medicine and Molecular Science, Graduate School of Medicine, Osaka University, 2-2 Yamadaoka, Suita, Osaka, 565-0871, Japan SO Arteriosclerosis Thrombosis and Vascular Biology, (June, 2000)

Vol. 20, No. 6, pp. 1595-1599. print.

ISSN: 1079-5642

LA English

ED Entered STN: 19 Jul 2000

AB Adiponectin is a novel, adipose-specific protein abundantly present in the circulation, and it has antiatherogenic properties. We Last Updated on STN: 7 Jan 2002

mass index (BMI)-matched nondiabetic and type 2 diabetic subjects with and 6.6+-0.4 mug/mL, P<0.001 in men; 6.3+-0.8 versus 7.6+-0.7 mug/mL in those in nondiabetic subjects (6.6+-0.4 versus 7.9+-0.5 mug/mL in men. 7.6+-0.7 versus 11.7+-1.0 mug/mL in women; P<0.001). The plasma</p> ower than those of diabetic patients without CAD (4.0+-0.4 versus adiponectin in the diabetic subjects without CAD were lower than analyzed the plasma adiponectin concentrations in age- and body adiponectin concentrations of diabetic patients with CAD were without coronary artery disease (CAD). Plasma levels of

(r=-0.18, P<0.01) and glucose (r=-0.26, P<0.001) levels. In multivariate between diabetic patients with and without CAD. The presence of diabetic patients. Significant, univariate, inverse correlations were microangiopathy did not affect the plasma adiponectin levels in observed between adiponectin levels and fasting plasma insulin analysis, plasma insulin did not independently affect the plasma adiponectin levels. BMI, serum triglyceride concentration, and women). In contrast, plasma levels of leptin did not differ

the presence of diabetes or CAD remained significantly related to plasma elevated plasma adiponectin levels in the diabetic subjects as adiponectin concentrations. Weight reduction significantly

decreased plasma adiponectin concentrations in diabetes may be

well as the nondiabetic subjects. These results suggest that the

an indicator of macroangiopathy.

L12 ANSWER 57 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN AN 2001:42054 CAPLUS << LOGINID::20070127>>

DN 134:54607

TI The influence of the genes expressed in adipose tissue on diseases

AU Takahashi, Masahiko; Funahashi, Tohru

CS Dep. Intern. Med. Mol. Sci., Grad. Sch. Med., Osaka Univ., Japan SO Horumon to Rinsho (2000), 48(12), 1055-1062

CODEN: HORIAE; ISSN: 0045-7167

PB Igaku no Sekaisha

DT Journal; General Review

pathogenesis of coronary artery diseases, diabetes mellitus, hypertension, LA Japanese AB A review with 26 refs., on the pathol. of visceral fat syndrome, genes AB A review with

and other common diseases. Structure, distribution, and pathophysiol. functions of adiponectin/apM 1 (adipose most abundant gene transcript 1), plasminogen activator inhibitor 1, TNFa,

leptin, PPARg, SREBP, and aquaporin adipose are discussed

L12 ANSWER 58 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

STS

AN 2000:490437 BIOSIS <<LOGINID::20070127>> DN PREV200000490558

T1 Molecular mechanism of obesity-related diseases: Importance of adipocytokines.

AU Matsuzawa, Yuji [Reprint author]; Funahashi, Tohru; Kuriyama, Hiroshi;

CS Department of Internal Medicine and Molecular Science, Graduate School of Kihara, Shini

Medicine, Osaka University 2-2 B-5, Yamadaoka, Suita, Osaka, 565-087

SO imura, Hiroo; Kasuga, Masato; Nakao, Kazuwa. Int. Congr. Ser. - Excerpta Med., (1999) pp. 37-43. International Congress Series; Common disease: Genetic and pathogenetic aspects of multifactorial diseases.

Publisher: Elsevier Science B.V., Sara Burgerhartstraat 25, 1000 AE,

Meeting Info.: Proceedings of the Uehara Memorial Foundation Symposium on Amsterdam, Netherlands. Series: International Congress Series.

CODEN: EXMDA4. ISSN: 0531-5131. ISBN: 0-444-50200-9 (cloth). Common Disease. Tokyo, Japan. June 30-July 02, 1999.

Conference; (Meeting) Book; (Book Chapter)

Conference; (Meeting Paper)

L12 ANSWER 52 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

STS

AN 2001:564208 BIOSIS <<LOGINID::20070127>> DN PREV200100564208

TI PPARgamma agonist and antagonist.

AU Kadowaki, Takashi [Reprint author]

Department of Metabolic Diseases, Graduate School of Medicine, University of Tokyo, Tokyo, 113-8655, Japan S

kadowaki-3im@h.u-tokyo.ac.jp

SO Folia Pharmacologica Japonica, (November, 2001) Vol. 118, No. 5,

pp. 321-326. print.

CODEN: NYKZAU. ISSN: 0015-5691

DT Article

Entered STN: 5 Dec 2001 LA Japanese ED Entered ST

Last Updated on STN: 25 Feb 2002

a retinoid X receptor (RXR). Supraphysiological activation of PPARgamma by thiazolidinediones can reduce insulin resistance and hyperglycemia in AB Peroxisome proliferator-activated receptor gamma (PPARgamma) is a ligand-activated transcription factor and functions as a heterodimer with type 2 diabetes, but these drugs can also

investigated whether functional antagonism toward PPARgamma/RXR could be cause weight gain. Quite unexpectedly, a moderate reduction of PPARgamma Ala polymorphism in human PPARgamma has been shown to prevent insulin activity observed in heterozygous PPARgamma-deficient mice or the Pro 12 resistance and obesity induced by a high-fat (HF) diet. We used to treat obesity and type 2

with an RXR antagonist or a PPARgamma antagonist decreases triglyceride TG) content in white adipose tissue, skeletal muscle and liver. These diabetes. We show herein that moderate reduction of PPARgamma adiponectin levels, which increases fatty acid combustion and inhibitors potentiate leptin's effects and stimulated

treatment of heterozygous PPARgamma-deficient mice with an RXR antagonist suggest that appropriate functional antagonism of PPARgamma/RXR may be a and insulin resistance. Paradoxically, severe reduction of PPARgamma by or a PPARgamma antagonist depletes white adipose tissue and markedly dissipation, which increases TG content in skeletal muscle and the liver, thereby leading to the re-emergence of insulin resistance. Our data energy dissipation, thereby ameliorating HF diet-induced obesity decreases leptin and adiponectin levels and energy

L12 ANSWER 53 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN

ogical approach to protection against obesity and related

diseases such as type 2 diabetes.

2001:279651 CAPLUS << LOGINID:: 20070127>>

134:250319

II Insulin resistance and visceral obesity

AU Nishida, Makoto, Funahashi, Tohtu CS Dep. Intern. Med. Mol. Sci., Grad. Sch. Med., Osaka Univ., Japan SO Horumon to Rinsho (2001), 49(3), 227-233

CODEN: HORIAE; ISSN: 0045-7167

PB Igaku no Sekaisha

DT Journal; General Review
LA Japanese
AB A review with 37 refs., on the clin. importance of visceral fat syndrome, pathophysiol. functions of adipocytokines, structure and functions of adiponectin, mechanism of the induction of insulin resistance by

visceral fat accumulation, action mechanisms of thiazolidine derivs. (PPARg agonists), roles of free fatty acids in insulin resistance,

and involvement of adipocytokines (TNFa, leptin, and adiponectin) in insulin resistance. L12 ANSWER 55 OF 60 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

STS

6

AN 2001:441526 BIOSIS <<LOGINID::20070127>>

DN PREV200100441526

II Replenishment of fat-derived hormone adiponectin reverses insulin resistance in lipoatrophic diabetes and type 2

Terauchi, Yasuo [Reprint author]; Kubota, Naoto [Reprint author]; Waki AU Yamauchi, Toshimasa [Reprint author]; Kamon, Junji [Reprint author] diabetes.

[Reprint author]; Tobe, Kazuyuki [Reprint author]; Yoda, Madoka [Reprint Hironori [Reprint author]; Mori, Yasumichi [Reprint author]; Hara, Kazuo author]; Tomita, Motowo [Reprint author]; Froguel, Philippe [Reprint [Reprint author]; Akanuma, Yasuo [Reprint author]; Kimura, Satoshi

author]; Kadowaki, Takashi [Reprint author]

CS Tokyo, Japan

SO Diabetes, (June, 2001) Vol. 50, No. Supplement 2, pp. A70.

Association. Philadelphia, Pennsylvania, USA. June 22-26, 2001. American Meeting Info.: 61st Scientific Sessions of the American Diabetes

CODEN: DIAEAZ. ISSN: 0012-1797. Diabetes Association.

Conference; (Meeting) Ы

Conference; Abstract; (Meeting Abstract)

ED Entered STN: 19 Sep 2001

Last Updated on STN: 22 Feb 2002

- Last Updated on STN: 10 Jan 2002 Entered STN: 15 Nov 2000 LA English ED Entered (
- L12 ANSWER 59 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN AN 1999:687090 CAPLUS <<LOGNID::20070127>>

 - 132:32236
- Cell biology of visceral fat

AU Hotta, Kikuko; Matsuzawa, Yuji

- Department of Internal Medicine and Molecular Science, Graduate School of Medicine, Osaka University, Yamadaoka, Suita-shi, Osaka, 565-0871, Japan Ś
 - SO Nihon Yukagakkaishi (1999), 48(10), 963-970
 - CODEN: NIYUFC; ISSN: 1341-8327
 - PB Nihon Yukagaku Gakkai
- Journal; General Review apanese
- A review with 49 refs. Adipose tissue is a source of passively stored excess energy. Adipose tissue has been found to secrete various biol inhibitor (PAI)-1 and tumor necrosis factor (TNF) a, which affect active adipocytokines such as leptin, plasminogen activator ΑB
- adipose to possibly be essential to glycerol metabolism in adipocytes. The water channel transports glycerol as well as water, suggesting aquaporin Aquaporin adipose is also highly expressed in adipose tissue. This new adipocytokines should facilitate clarification of the mechanism for the iomeostasis throughout the body. Plasma adipocytokines increase in obesity and the accumulation of fat, especially visceral fat, may serve search for genes expressed in adipose tissue, novel adipose-specific present study. Adiponectin had a collagen-like sequence and was secreted into blood. Plasma adiponectin paradoxically decreased obesity, through enhanced secretion of the above compds. In genes, adiponectin and aquaporin adipose were isolated in the finding of genes specifically expressed in visceral fat and new to create greater insulin resistance or thrombotic tendency in development and complications of visceral fat accumulation. in obesity, but was expressed exclusively in adipose tissue.
- L12 ANSWER 60 OF 60 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 2000:101167 CAPLUS <<LOGIND::20070127>> DN 133:28940
- TI Molecular mechanism of obesity-related diseases: importance of adipocytokines
- Department of Internal Medicine and Molecular Science, Graduate School of Matsuzawa, Yuji; Funahashi, Tohru; Kuriyama, Hiroshi; Kihara, Shinji ΑŪ S
 - International Congress Series (1999), 1181(Common Disease: Medicine, Osaka University, Suita, 565-0871, Japan S
- Genetic and Pathogenetic Aspects of Multifactorial Diseases), 37-43

CODEN: EXMDA4; ISSN: 0531-5131

- PB Elsevier Science B.V.
- DT Journal; General Review
- human disorders including diabetes mellitus, hyperlipidemia, hypertension large-scale-random sequencing and revealed that adipose tissue, especially apprx.30% and .apprx.20% of the total genes in visceral and s.c. adipose. related to atherogenesis such as plasminogen activator inhibitor-1 (PAI-1) and heparin-binding EGF-like growth factor (HB-EGF) were found in the Plasma levels of PAI-1 were closely correlated with visceral adiposity in adipose tissue, a systemic anal. of expressed genes was performed using visceral adipose tissue expressed numerous genes for secretory proteins ibrary. PAI-1, a regulator of fibrinolytic system, was overexpressed in called "adipocytokines" which affect biol. function of each target organ. have revealed that adipose tissue is not simply an energy storage organ but also an endocrine organ secreting a variety of bioactive substances appetite and energy expenditure. To clarify the mol. characteristics of and atherosclerotic vascular disease. Recent studies on adipocyte biol tissue, resp.). Among these secretory proteins, active genes reputedly famous adipocytokines, which have an important role in controlling For example, TNF-a from adipose tissues is one of the key factors LA English
 AB A review, with 12 refs. Obesity is a major cause of common AB. for the development of insulin resistance. Leptin is another the visceral adipose tissue in an animal model of obesity.

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS obesity-related diseases including diabetes mellitus and vascular RECORD

substances, adipocytokines. Hypersecretion of adipocytokines such as

adiponectin in obese state may relate to the pathogenesis of

PAI-1 or TNF-a and hyposecretion of those such as

proliferation and inhibition of adhesion mol. expression in endothelial

anti-atherogenic property such as inhibition of smooth muscle cell

adiponectin was found. This novel mol. is suggested to have an

eptin were neg. correlated with body mass indexes. Thus, adipose

cells, etc. The plasma levels of adiponectin unlike

tissue acts as an endocrine organ secreting a variety of bioactive

human subjects. A novel adipose-specific collagen-like mol. named

ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> D bib ABS L149, 14

L14 ANSWER 9 OF 15 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on

AN 2002:266783 BIOSIS <<LOGINID::20070127>>

- The mechanisms by which PPARgamma regulates insulin sensitivity.
- AU Yamauchi, Toshimasa [Reprint author]; Kadowaki, Takashi [Reprint author] Dept. of Metabolic Disease, Graduate Sch. of Med., Univ. of Tokyo, Tokyo, S
- I, pp. 24P. print.

SO Japanese Journal of Pharmacology, (2002) Vol. 88, No. Supplement

- Meeting Info:: 75th Annual Meeting of the Japanese Pharmacological Society. Kumamoto, Japan. March 13-15, 2002. Japanese Pharmacological
- CODEN: JJPAAZ, ISSN: 0021-5198.
- Conference; Abstract; (Meeting Abstract) DT Conference; (Meeting)
 - LA English ED Entered STN: 1 May 2002
- Last Updated on STN: 1 May 2002

L14 ANSWER 14 OF 15 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

STS

등

- AN 2001:187493 BIOSIS <<LOGINID::20070127>> DN PREV200100187493
- TI Leptin signals and secretions from white adipose tissue.
- AU Trayhum, Paul [Reprint author]; Beattie, John H. [Reprint author]; Rayner, D. Vernon [Reprint author]
 CS Rowett Research Institute, Bucksburn, Aberdeen, AB21 9SB, UK
- SO Heldmaier, Gerhard; Klingenspor, Martin. (2000) pp. 459-469.
- Publisher: Springer-Verlag GmbH and Co. KG, Heidelberger Platz 3, D-14197, Life in the cold. print.
 - Berlin, Germany; Springer-Verlag New York Inc., 175 Fifth Avenue, New York, NY, 10010-7858, USA.
 - Meeting Info.: Eleventh International Hibernation Symposium. Jungholz, Austria. August 13-18, 2000
 - SBN: 3-540-67410-1 (cloth)
 - Book
- Conference; (Meeting)
- Book; (Book Chapter)
- Conference; (Meeting Paper)
 - English ۲
- Last Updated on STN: 18 Feb 2002 Entered STN: 20 Apr 2001
- => D bib Abs L16 1, 3, 10, 14, 30, 39, 42, 43, 50, 65, 73-75,91, 126, 157, 167, 178, 183

- L16 ANSWER I OF 191 MEDLINE on STN
- 2002669523 MEDLINE <<LOGINID::20070127>> AN 2002669523 MEDLIT DN PubMed ID: 12429885

- TI Resistin and adiponectin—of mice and men.
 AU Sturnvoll Michael; Haring Hans
 SO Obesity research, (2002 Nov) Vol. 10, No. 11, pp. 1197-9. Ref: 28
- Journal code: 9305691. ISSN: 1071-7323
- United States Editorial Ç
- General Review; (REVIEW)
 - LA English
- FS Priority Journals EM 200305
- ED Entered STN: 14 Nov 2002
- Last Updated on STN: 14 May 2003 Entered Medline: 13 May 2003

L16 ANSWER 3 OF 191 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

6

- AN 2003:32810 BIOSIS <<LOGINID::20070127>>
 - DN PREV200300032810
- II Increased fat intake, impaired fat oxidation, and failure of fat cell proliferation result in ectopic fat storage, insulin resistance, and
 - type 2 diabetes mellitus.
- AU Ravussin, Eric [Reprint Author]; Smith, Steven R.
- CS Pennington Biomedical Research Center, 6400 Perkins Road, Baton Rouge, LA. 70808-4124, USA

 - SO Klimes, Iwar [Editor, Reprint Author]; Sebokova, Elena [Editor]; Howard, ravusse@pbrc.edu
- Barbara V. [Editor]; Ravussin, Eric [Editor]. (2002) pp. 363-378. Lipids and insulin resistance: The role of fatty acid metabolism
- Publisher: New York Academy of Sciences, 2 East 63rd Street, New York, NY, and fuel partitioning. print.
 - Meeting Info.: Fourth International Smolenice Insulin Symposium on Lipids and Insulin Resistance: The Role of Fatty Acid Metabolism and Fuel 10021, USA. Series: Annals of the New York Academy of Sciences.
- ISSN: 0077-8923 (ISSN print). ISBN: 1-57331-368-8 (cloth), 1-57331-369-6 Partitioning. Smolenice, Slovakia. August 29-September 02, 2001
- DT Book; (Book Chapter)
- Conference; (Meeting)
- Conference; (Meeting Paper)
 - LA English ED Entered STN: 8 Jan 2003

Last Updated on STN: 8 Jan 2003

L16 ANSWER 10 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:775209 CAPLUS <<LOGIND::20070127>>

DN 138:37209

TI Removal of visceral fat prevents insulin resistance and glucose

intolerance of aging: an adipokine-mediated process?

AU Gabriely, Ilan; Ma, Xiao Hui; Yang, Xiao Man; Arzmon, Gil; Rajala, Michael

CS Diabetes Research and Training Center and Division of Endocrinology, W.; Berg, Anders H.; Scherer, Phillip; Rossetti, Luciano; Barzilai, Nir

Department of Medicine, Institute for Aging Research, Albert Einstein College of Medicine, Bronx, NY, 10461, USA

SO Diabetes (2002), 51(10), 2951-2958

CODEN: DIAEAZ; ISSN: 0012-1797

PB American Diabetes Association

LA English Journal

AB Age-dependent changes in insulin action and body fat distribution are risk factors for the development of type 2 diabetes

To examine whether the accumulation of visceral fat (VF) could play a direct role in the pathophysiol. of insulin resistance and type

2 diabetes, we monitored insulin action, glucose

removal of VF in aging (20-mo-old) F344/Brown Norway (FBN) and in Zucker tolerance, and the expression of adipo-derived peptides after surgical

action were markedly impaired in aging FBN rats, and extraction of VF (accounting for apprx.18% of their total body fat) was sufficient to Diabetic Fatty (ZDF) rats. As expected, peripheral and hepatic insulin restore peripheral and hepatic insulin action to the levels of young rats.

When examined at the mechanistic level, removal of VF in ZDF rats prevented the progressive decrease in insulin action and delayed the onset of

s.c. (SC) adipose tissue were markedly decreased after VF removal (by diabetes, but VF extraction did not alter plasma free fatty acid levels. approx. three- and twofold, resp.). Finally, extracted VF retained However, the expression of tumor necrosis factor-a and leptin in

accumulation of VF. This study documents a cause-and-effect relationship apprx.15-fold higher resistin mRNA compared with SC fat. Our data suggest that insulin resistance and the development of diabetes can be significantly reduced in aging rats by preventing the age-dependent between VF and major components of the metabolic syndrome.

RE.CNT 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 14 OF 191 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

DUPLICATE 8

5

AN 2002:416819 BIOSIS <<LOGINID::20070127>> DN PREV200200416819

2 diabetes: A candidate gene for the insulin resistance II Association of adiponectin mutation with type

Kondo, Hidehiko; Shimomura, Iichiro; Matsukawa, Yuko; Kumada, Masahiro; Kawamoto, Toshiharu; Sumitsuji, Satoru; Funahashi, Tohru [Reprint author]; Takahashi, Masahiko; Matsuda, Morihiro; Ouchi, Noriyuki; Kihara, Shinji; Matsuzawa, Yuji ΑŪ

CS Department of Internal Medicine and Molecular Science, Osaka University Graduate School of Medicine, B5 2-2, Yamadaoka, Suita, Osaka, 565-0871 Japan

tohru@imed2.med.osaka-u.ac.jp SO Diabetes, (July, 2002) Vol. 51, No. 7, pp. 2325-2328. print. CODEN: DIAEAZ. ISSN: 0012-1797.

Article

ED Entered STN: 31 Jul 2002 LA English

Last Updated on STN: 31 Jul 2002

protein contains a collagen-like domain and a C1q-like globular domain. protease-generated globular segment enhances fatty acid oxidation in muscles, thereby modulating lipid and glucose metabolism. Plasma protein produced and secreted exclusively from adipose tissue. The AB Adiponectin, also referred to as AdipoQ or ACRP30, is a plasma insulin resistance. A recent genome-wide scan study mapped a adiponectin levels are inversely correlated with the severity of

the metabolic syndrome to chromosome 3q27, where the adiponectin gene is located. Here, we screened Japanese patients with type susceptibility locus for type 2 diabetes and

missense mutations (R112C, 1164T, R221S, and H241P) in the globular domain. Among these mutations, the frequency of 1164T mutation was subjects for mutations in adiponectin gene. We identified four 2 diabetes and age- and BMI-matched nondiabetic control

were lower than those of subjects without the mutation. All the subjects carrying 1164T mutation showed some feature of metabolic syndrome, BMI-matched control subjects (P < 0.01). Furthermore, plasma adiponectin concentrations of subjects carrying 1164T mutation significantly higher in type 2 diabetic patients than in age- and

Our findings suggest that 1164T mutation is associated with low plasma including hypertension, hyperlipidemia, diabetes, and atherosclerosis. adiponectin concentration and type 2

L16 ANSWER 30 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN DUPLICATE

AN 2003:275394 CAPLUS << LOGINID::20070127>>

- DN 138:399418
- TI Adiponectin a link between obesity, atherosclerosis and diabetes
- Sieminska, Lucyna; Marek, Bogdan; Kajdaniuk, Dariusz; Kos-Kudla, Beata; Czemecka, Dagmara
 - Zakl. Patofizjol., Katedra Patofizjol. i Endokrynol., Sl. Akad. Medyczna, Zabrze, Pol. S
 - SO Polskie Archiwum Medycyny Wewnetrznej (2002), 108(6), 1245-1251
 - CODEN: PAMWAL; ISSN: 0032-3772
 - PB Wydawnictwo Medyczne Urban & Partner DT Journal; General Review
- AB A review. The topics include the biochem. of adiponectins, their cellular physiol., and possible roles in pathogenesis of obesity, atherosclerosis, and diabetes mellitus in humans
- LI6 ANSWER 39 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN DUPLICATE
- AN 2003:65783 CAPLUS <<LOGINID::20070127>>
 - DN 138:265785
- TI Adipose tissue hormones

AU Guerre-Millo, M.

- CS Centre de Recherche des Cordeliers, Universite Pierre et Marie Curie, Paris, 75006, Fr.
 - SO Journal of Endocrinological Investigation (2002), 25(10),
 - 855-861
 - CODEN: JEIND7; ISSN: 0391-4097
- PB Editrice Kurtis s.r.l.
 DT Journal; General Re
- Journal; General Review English ۲
- secretes a number of peptide hormones, including leptin, several cytokines, AB A review. It is now widely accepted that white adipose tissue (WAT) longer considered only an energy storage tissue but a major endocrine secretory function has shifted the authors' view of WAT, which is no adipsin and acylation-stimulating protein (ASP), angiotensinogen, plasminogen activator inhibitor-1 (PAI-1), adiponectin, resistin etc., and also produces steroids hormones. This newly discovered
- glucose and lipid metabolism, vascular homeostasis, immune response and even reproduction Virtually all known adipose secreted proteins are dysregulated secretion by focusing on protein and steroid hormones. Regulation of WAT when the WAT mass is markedly altered, either increased in the obese state organ, at the heart of a complex network influencing energy homeostasis, or decreased in lipoatrophy. This strongly implicates adiposesecreted and cachexia. This review discusses the physiol, relevance of adipose secretion by the major regulatory factors impinging on the adipocytes, products in the ethiopathol. and/or complications of both obesity

- (TZD) will be addressed. The rationale for therapeutic strategies aimed at compensating adverse effects resulting from overprodn. or lack of a specific adipose secretory product will be discussed
- RE.CNT 53 THERE ARE 53 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L16 ANSWER 42 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:453341 CAPLUS <<LOGINID::20070127>>
 - - 137:61015

- TI Obesity and adipo-science
 AU Yamauchi, Toshimasa; Kadowaki, Takashi
 CS Grad. Sch. Med., The Univ. Tokyo, Japan
 SO Rinsho Eiyo (2002), 100(6, Rinjizokango), 745-750 CODEN: RNEYAW; ISSN: 0485-1412

- PB Ishiyaku Shuppan
 DT Journal; General Review
 LA Japanese
 AB A review on improvement of insulin resistance by PPARg agonists,
 - thiazolidine derivs, via acceleration of adipocyte differentiation and apoptosis, functions of PPARg as a thrift gene, PPARg2 gene
 - polymorphism and type 2 diabetes mellitus, treatment of type 2 diabetes mellitus by
- PPARg inhibitors, regulation of insulin sensitivity by PPARg, and role of adiponectin in regulation of insulin sensitivity.
- L16 ANSWER 43 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:588796 CAPLUS <<LOGIND::20070127>> DN 138:130470 TI Tailor-made medicine for obesity
- AU Hotta, Kikuo CS SNP Research Center, Institute of Physical and Chemical Research, Japan SO Igaku no Ayumi (2002), 201(9), 725-728 Jgaku no Ayumi (2002), 201(9), 725-728CODEN: IGAYAY; ISSN: 0039-2359
 - - PB Ishiyaku Shuppan
- DT Journal; General Review
- LA Japanese AB A review, discussing the development of tailor-made medicine for obesity with regards to genomes and SNP related to
 - adiponectins and adipocytokines.
- L16 ANSWER 50 OF 191 EMBASE COPYRIGHT (c) 2007 Elsevier B.V. All rights reserved on STN DUPLICATE 23
 - AN 2002397931 EMBASE <<LOGINID::20070127>>
 - TI Diabetes, obesity, and Acrp30/adiponectin.

AU Hug C.; Lodish H.F.

CS Dr. H.F. Lodish, Whitehead Inst. for Biomed. Research, 9 Cambridge Center, Cambridge, MA 02142, United States. lodish@wi.mit.edu

BioTechniques, (2002) Vol. 33, No. 3, pp. 654-662. S

Refs: 33

SSN: 0736-6205 CODEN: BTNQDO

United States ح

Journal; General Review

Clinical Biochemistry Endocrinology 83

English 5

ED Entered STN: 2 Dec 2002

Last Updated on STN: 2 Dec 2002

DATA NOT AVAILABLE FOR THIS ACCESSION NUMBER

L16 ANSWER 65 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN DUPLICATE

AN 2002:832209 CAPLUS <<LOGINID::20070127>>

DN 138:117693

Adiponectin - its role in metabolism and beyond

Stefan, N.; Stumvoll, M.

Clinical Nutrition and Metabolism Section, NIDDK, NIH, Phoenix, AZ, 85016,

SO Hormone and Metabolic Research (2002), 34(9), 469-474 CODEN: HMMRA2; ISSN: 0018-5043

Georg Thieme Verlag В

Journal; General Review

English

increased by insulin like growth factor-1 and ionomycin, and decreased by itributable to enhanced suppression of glucose production, but beneficial issue-derived protein (adipocytokine) with important metabolic effects. increases insulin sensitivity and improves glucose tolerance in various and cAMP. Data for insulin are somewhat inconclusive. Moreover, nay contribute to the decrease in whole-body insulin sensitivity that unimal models. This insulin-sensitizing effect appears to be mostly thousand times; they decrease with obesity and are pos. associated of peroxisome proliferator-activated receptor (PPAR)-g. Besides t is exclusively expressed in adipose tissue and released into the adiponectin expression and secretion are increased by activators with whole-body insulin sensitivity. Therefore, low adiponectin tumor necrosis factor-a, glucocorticoids, b-adrenergic agonists adiponectin concus. exceed those of any other hormone by a inhibiting inflammatory pathways, recombinant adiponectin effects on muscle cannot be excluded. In humans, plasma AB A review. Adiponectin is a recently identified adipose circulation. Adiponectin expression and/or secretion is

PPAR-g - may be associated with hypoadiponectinemia, insulin resistance accompanies obesity. Furthermore, there is increasing evidence that genetic variants in the adiponectin gene itself and/or in adiponectin may reflect PPAR-g activity in vivo. Finally, genes encoding adiponectin-regulatory proteins - such as and type 2 diabetes. This suggests that

development of drugs improving insulin sensitivity and glucose tolerance.

RECNT 58 THERE ARE 58 CITED REFERENCES AVAILABLE FOR THIS

reversal or alleviation of hypoadiponectinemia may represent a target for

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 73 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN DUPLICATE

AN 2002:587011 CAPLUS <<LOGINID::20070127>> DN 137:38287 TI Increased fat intake, impaired fat oxidation, and failure of fat cell proliferation result in ectopic fat storage, insulin resistance, and

type 2 diabetes mellitus

Ravussin, Eric; Smith, Steven R.

Pennington Biomedical Research Center, Baton Rouge, LA, 70808-4124, USA Annals of the New York Academy of Sciences (2002), 967(Lipids

and Insulin Resistance), 363-378

CODEN: ANYAA9; ISSN: 0077-8923 PB New York Academy of Sciences DT Journal; General Review

LA English

AB A review. It is widely accepted that increasing adiposity is associated with insulin resistance and increased risk of type 2

hypothesis. As such, two new paradigms have emerged that may explain the established links between adiposity and disease. Three lines of evidence probably the beta cell. The importance of this finding is exemplified by Third, increased fat cell size is highly associated with insulin resistance the portal/visceral hypothesis. This hypothesis proposes that increased lipodystrophy, results in severe insulin resistance and diabetes. This is adequate adipose tissue mass in either mice or humans, also known as several studies demonstrating that the degree of lipid infiltration into thought to be the result of ectopic storage of lipid into liver, skeletal muscle, and the pancreatic insulin-secreting beta cell. Second, most obese patients also shunt lipid into the skeletal muscle, the liver, and fatty acid flux and inhibition of insulin action via Randle's effect in adiposity, particularly in the visceral depots, leads to increased free support the ectopic fat storage syndrome. First, failure to develop insulin-sensitive tissues. Recent data do not entirely support this skeletal muscle and liver correlates highly with insulin resistance diabetes. The predominant paradigm used to explain this link is

gland, secreting numerous factors with potent effects on the metabolism of distant tissues. These two new paradigms provide a framework to advance our understanding of the pathophysiol. of the insulin-resistance syndrome. RE.CNT 118 THERE ARE 118 CITED REFERENCES AVAILABLE FOR THIS and insulin resistance. The endocrine paradigm developed in parallel with support the acquired lipodystrophy hypothesis as a link between adiposity the failure of the adipose tissue mass to expand and thus to accommodate variety of endocrine hormones, such as leptin, interleukin-6, angiotensin From this viewpoint, adipose tissue plays a critical role as an endocrine the ectopic fat storage syndrome hypothesis. Adipose tissue secretes a and the development of diabetes. Increased fat cell size may represent an increased energy influx. Taken together, these three observations adiponectin (also called ACRP30 and adipoQ), and resistin.

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L16 ANSWER 74 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN
 - AN 2002:578380 CAPLUS <<LOGINID::20070127>>
 - DN 138:87270
- II Adiponectin antidiabetic and antiatherogenic adipocytokine
- Shimomura, lichiro; Hunahashi, Toru; Kihara, Shinji; Matsuzawa, Yuji Dep. Internal Med. Molecular Sci., Grad. Sch. Med., Osaka Univ., Suita, ပ္ပ
 - SO Naibunpi, Tonyobyoka (2002), 14(4), 361-366 CODEN: NATOFF, ISSN: 1341-3724 565-0871, Japan
- PB Kagaku Hyoronsha
- Journal; General Review Japanese
- AB A review, on the concept of adipocytokines, especially adiponectin, in adipocyte-specific hormones; and pathophysiol. roles of ipodystrophy and obesity; adiponectin as

adiponectin in diabetes, atherosclerosis, and metabolic disorders.

- 16 ANSWER 75 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN DUPLICATE
- AN 2002:907355 CAPLUS <<LOGINID::20070127>>
 - DN 138:300852
- TI An adipocentric view of signaling and intracellular trafficking
- Mora, Silvia; Pessin, Jeffrey E.
- Department of Physiology and Biophysics, The University of Iowa, Iowa, IA, USA S
- SO Diabetes/Metabolism Research and Reviews (2002), 18(5), 345-356 CODEN: DMRRFM; ISSN: 1520-7552
 - PB John Wiley & Sons Ltd.
- Journal; General Review

stimulate both glucose uptake and lipogenesis. Conventional wisdom held involved in fatty acid and glucose metabolism (leptin, Acrp30, resistin and site for whole body energy storage mainly in the form of triglycerides and acylation stimulation protein). Despite the large number of mols. secreted models and humans also leads to metabolic disorders that result in severe opposite functions can be resolved by the establishment of adipocytes not regulating insulin secretion, insulin action, glucose and lipid metabolism, secretory factors include enzymes (lipoprotein lipase (LPL) and adipsin), poorly understood. In this article, we will review the current knowledge increased adipose tissue mass and/or increased lipolysis and circulating mol. function of these adipocyte-derived signals are poorly understood, growth factors [vascular endothelial growth factor (VEGF)], cytokines secretes a variety of signaling mols. into the circulation. Although the of the trafficking and secretion processes that take place in adipocytes, focusing our attention on two of the best characterized adipokine mols resistance and perhaps diabetes. However, it has become increasingly energy balance, host defense and reproduction The diversity of these studied regulated membrane proteins, the GLUT4 glucose transporter only as a fuel storage depot but also as a critical endocrine organ that they play a central role in the maintenance of energy homeostasis by apparent that loss of adipose tissue (lipodystrophies) in both animal states of insulin resistance and potential diabetes. These apparently (turnor necrosis factor-a, interleukin 6) and several other hormones by adipocytes, our understanding of the pathways and mechanisms controlling intracellular trafficking and exocytosis in adipocytes is fatty acids. This occurs through the ability of insulin to markedly that defects in fuel partitioning into adipocytes either because of (leptin and adiponectin) and on one of the most intensively free fatty acids resulted in dyslipidemia, obesity, insulin

RE.CNT 174 THERE ARE 174 CITED REFERENCES AVAILABLE FOR THIS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L16 ANSWER 91 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN AN 2002:378204 CAPLUS <<LOGINID::20070127>> DN 137:308016 TI The role of adiponectin in obesity, insulin
- resistance, and type 2 diabetes. The
- fat-derived hormone adiponectin reverses insulin resistance
 - AU Yamauchi, Toshimasa; Kadowaki, Takash associated with both lipoatrophy and obesity
- CS Department of Metabolic Diseases, Graduate School of Medicine, University
 - SO Naibunpi, Tonyobyoka (2002), 14(2), 172-179 of Tokyo, Tokyo, 113-8655, Japan
 - CODEN: NATOFF; ISSN: 1341-3724

- Kagaku Hyoronsha
- Journal; General Review
 - LA Japanese
- AB A review, on roles of adiponectin in regulation of insulin sensitivity, adiponectin gene as the major disease-sensitive sensitivity, discussing adiponectin expression and insulin hormone derived from white adipocytes; and adiponectin diabetes and adiponectin supplement in improvement of diabetes; adiponectin as the major insulin sensitive deficiency induction of obesity and type 2 gene in Japanese population with type 2
- L16 ANSWER 126 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN
 - AN 2002:16380 CAPLUS <<LOGINID::20070127>>

insulin resistance

- lat-derived hormone adiponectin reverses insulin resistance TI The role of adiponectin in obesity, insulin resistance, and type 2 diabetes: The
 - associated with both lipoatrophy and obesity
- AU Yamauchi, Toshimasa; Kadowaki, Takashi
- CS Department of Metabolic Diseases, Graduate School of Medicine, University of Tokyo, Japan
 - SO Jikken Igaku (2001), 19(17), 2301-2305
 - CODEN: JIIGEF; ISSN: 0288-5514
 - PB Yodosha
- Journal; General Review
- AB A review discussing increased adiponectin expression in Japanese

heterozygous PPARg deficiency with improved insulin sensitivity; genetic variations in the adiponectin gene associated with Japanese population; fat-derived adiponectin as insulin increased risk of type 2 diabetes in

sensitive hormone; and insulin resistance induced by adiponectin

deficiency.

L16 ANSWER 157 OF 191 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation 6

SIS

AN 2001:441526 BIOSIS <<LOGINID::20070127>>
DN PREV200100441526
TI Replenishment of fat-derived hormone adiponectin reverses insulin resistance in lipoatrophic diabetes and type 2 diabetes.

AU Yamauchi, Toshimasa [Reprint author]; Kamon, Junji [Reprint author]; Terauchi, Yasuo [Reprint author]; Kubota, Naoto [Reprint author]; Waki,

[Reprint author]; Tobe, Kazuyuki [Reprint author]; Yoda, Madoka [Reprint author]; Tomita, Motowo [Reprint author]; Froguel, Philippe [Reprint Hironori [Reprint author]; Mori, Yasumichi [Reprint author]; Hara, Kazuo [Reprint author]; Kimura, Satoshi author]; Kadowaki, Takashi [Reprint author]

- CS Tokyo, Japan SO Diabetes, (June, 2001) Vol. 50, No. Supplement 2, pp. A70.

Association. Philadelphia, Pennsylvania, USA. June 22-26, 2001. American Meeting Info.: 61st Scientific Sessions of the American Diabetes Diabetes Association.

CODEN: DIAEAZ. ISSN: 0012-1797.

- DT Conference; (Meeting)
- Conference; Abstract; (Meeting Abstract)
 - LA English ED Entered STN: 19 Sep 2001
- Last Updated on STN: 22 Feb 2002
- L16 ANSWER 167 OF 191 CAPLUS COPYRIGHT 2007 ACS on STN
 - AN 2000:777806 CAPLUS <<LOGINID::20070127>>
 - II Adipocytokines DN 133:308338
- AU Yokota, Takafumi; Takahashi, Masahiko; Funahashi, Tohru CS Grad. Sch. Med., Osaka Univ., Japan SO Ensho to Men'eki (2000), 8(6), 624-629
- CODEN: ENMEFA; ISSN: 0918-8371
- DT Journal; General Review

PB Sentan Igakusha

- LA Japanese A review with 11 refs., on the expression of adipocytokines in visceral fat and their involvement in obesity complications, enhanced expression of PAI-1 in visceral fat, and structure and pathophysiol. functions of adiponectin. The decrease of adiponectin

and suppression of the proliferation and functions of macrophages by expression in humans with obesity and coronary artery diseases, adiponectin are discussed.

L16 ANSWER 178 OF 191 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation

STS

AN 2000:490437 BIOSIS <<LOGINID::20070127>> DN PREV200000490558

adipocytokines.

II Molecular mechanism of obesity-related diseases: Importance of

AU Matsuzawa, Yuji [Reprint author]; Funahashi, Tohru; Kuriyama, Hiroshi;

Kihara, Shinji

- CS Department of Internal Medicine and Molecular Science, Graduate School of Medicine, Osaka University 2-2 B-5, Yamadaoka, Suita, Osaka, 565-0871
- SO Imura, Hiroo; Kasuga, Masato; Nakao, Kazuwa. Int. Congr. Ser. Excerpta Med., (1999) pp. 37-43. International Congress Series; Common disease: Genetic and pathogenetic aspects of multifactorial diseases.

Publisher: Elsevier Science B.V., Sara Burgerhartstraat 25, 1000 AE, Amsterdam, Netherlands. Series: International Congress Series.

Meeting Info.: Proceedings of the Uehara Memorial Foundation Symposium on Common Disease. Tokyo, Japan. June 30-July 02, 1999.

CODEN: EXMDA4. ISSN: 0531-5131. ISBN: 0-444-50200-9 (cloth)

DT Book

Conference; (Meeting)

Conference; (Meeting Paper) Book; (Book Chapter)

English ۲

Last Updated on STN: 10 Jan 2002 Entered STN: 15 Nov 2000

L16 ANSWER 183 OF 191 MEDLINE on STN

AN 1999240218 MEDLINE << LOGINID::20070127>>

DN PubMed ID: 10225688

obesity.

Tl Role of adipocytokines on the pathogenesis of atherosclerosis in visceral

- AU Funahashi T; Nakamura T; Shimomura I; Maeda K; Kuriyama H; Takahashi M; Arita Y; Kihara S; Matsuzawa Y
 - CS The Second Department of Internal Medicine, Osaka University Medical School, Suita.
 - SO Internal medicine (Tokyo, Japan), (1999 Feb) Vol. 38, No. 2, pp.

202-6. Ref: 14

Journal code: 9204241. ISSN: 0918-2918. CY Japan

DT Journal; Article; (JOURNAL ARTICLE) General Review; (REVIEW)

LA English FS Priority Journals EM 199906

Last Updated on STN: 12 Jul 1999 Entered Medline: 18 Jun 1999 ED Entered STN: 12 Jul 1999

a major cause of atherosclerotic vascular disease in industrial countries. adipose tissue is not simply an energy storage organ but it also secretes a variety of molecules which affect the metabolism of the whole body. Recent advances in the biology of adipose tissue have revealed that AB Obesity which is defined as accumulation of excess body fat, is

that adipose tissue, especially visceral fat expressed numerous genes for plasminogen activator-1 (PAI-1), which is a regulator of the fibrinolytic Through a systematic search of active genes in adipose tissue, we found visceral fat adiposity. Thus, PAI-1 secreted from visceral fat may play secretory proteins (about 30% of total genes analyzed). Among them, system, was overexpressed in the visceral fat in an animal model of Adiponectin, a novel adipose-specific gene product, which has a Dysregulated secretion of adiponectin may be related to vascular adipose tissue (adipocytokines) may have important roles in the disease in obesity. Biologically active molecules secreted from matrix-like structure, is abundantly present in the bloodstream. obesity. Plasma levels of PAI-1 were closely correlated with some role in thrombotic vascular disease in visceral obesity. development of atherosclerotic disease in obesity.